Solving the structure of disordered mixed salts

A. Frenkel
School of Physics and Astronomy, Raymond and Beverly Sackler Faculty of Exact Sciences, Tel Aviv University, Ramat Aviv 69978, Israel

E. A. Stern
Department of Physics FM-15, University of Washington, Seattle, Washington 98195

A. Voronel
School of Physics and Astronomy, Raymond and Beverly Sackler Faculty of Exact Sciences, Tel Aviv University, Ramat Aviv 69978, Israel

M. Qian
Princeton Materials Institute, Princeton, New Jersey 08540

M. Newville
Department of Physics FM-15, University of Washington, Seattle, Washington 98195
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A detailed x-ray-absorption fine-structure (XAFS) investigation of two mixed alkali halides Rb$_{0.75}$K$_{0.25}$Br and RbBr$_{0.42}$Cl$_{0.58}$ was performed. The concentrations of the mixtures had been chosen to produce a single homogeneous phase for each, and it was checked by XAFS that the salts were randomly mixed on the atomic level. Detailed analysis of the data including multiple-scattering contributions revealed an rms buckling angular deviation of both mixtures from the average NaCl collinear structure of 7°-9°. The angles are defined by three atomic positions determined through double- and triple-scattering paths. These angles are new parameters which should be added to characterize the buckled crystals. Adding to diffraction results the parameters determined from XAFS as input into a molecular-dynamics simulation the structures of the mixed salts with their fluctuations about the NaCl structure are solved and displayed.

I. INTRODUCTION

The structure of solids is determined mainly by diffraction. For pure elements and compounds where the atoms are arranged in an ordered array, the sharp diffraction Bragg peaks give enough information to solve the structure. When elements and/or compounds are alloyed or mixed together, producing a single phase, the resultant solid still generally diffracts Bragg peaks, but in this case the Bragg reflections determine only the average structure, leaving unsolved the actual atomic deviation about this average. Typically the average structure has lattice constants which approximately vary with composition according to the virtual crystal approximation (VCA). The VCA states that the lattice constants approximately satisfy the Vegard's law of a linear interpolation between the corresponding pure components lattice constants, $a_1$ and $a_2$:

$$a_{\text{mix}} = xa_1 + (1-x)a_2.$$

(1)

The Bragg diffraction in such mixtures indicates that the average structure of these materials is still periodic. In this paper we address the question of determining the local structure and the specific character of its deviation from the average, which gives a complete solution to the structure. The method of how this is done using the x-ray-absorption fine-structure (XAFS) technique has previously been outlined, and in this paper we give further details and more results.

XAFS measurements in general imply that, locally, the atoms do not reside on the periodic structure. Indeed, it has been shown by XAFS in a variety of cases that the actual nearest neighbor distances in alloys and mixed salts are different from the average distances calculated from the lattice constant assuming a perfect periodic lattice. Thus it is clear that, generally speaking, the atoms are not sitting on their lattice sites of the average structure, but it has never been defined what is the actual structure locally and how to determine it quantitatively. In this paper our goal is to define in detail and to visualize the local structure of the two analogous systems of mixed salts, Rb$_x$K$_{1-x}$Br and RbBr$_x$Cl$_{1-x}$, by thorough analysis of their XAFS data. This choice of samples is not incidental. While the crystalline symmetries of these two mixtures are similar and the size difference between randomized K and Br ions is roughly equal to that between Cl and Br ($\approx 0.15\,\text{Å}$), and consequently their phase diagrams are similar, the absolute sizes of the randomized atoms are quite different. Since the randomized atoms also have different charges, it is not obvious...
that their local structure deviations will be similar. 

The knowledge of the actual locations of atoms in disordered solids and the character of their deviation from the average structure is necessary for detailed understanding of their properties. The dielectric, optical, electrical as well as mechanical properties of single-phase mixed salts are expected to depend on the character of distortion of the perfect structure. It has recently been pointed out that also the melting of such disorderly alloyed solids depends on deviation from the perfect structure. The larger is the initial deviation from the perfect structure, the more it depresses the melting point. 

In our analysis of XAFS data we utilize the theoretical program FEPF which includes multiple-scattering contributions to XAFS on a path by path basis. In particular we are able to estimate three-body correlations, which go beyond the usual two-body correlations normally taken into account by other means. The accuracy and power of FEPF have been demonstrated in describing the structure of the pure salts RbBr, RbCl, KBr, constituting the elements of both of the series of mixtures chosen for our study. The reliability of FEPF has also been proven by its determining various distortions in several perovskites in close agreement with diffraction results.

An outline of the paper is as follows. In Sec. II the experimental details are presented. The data analysis is discussed in Sec. III. Section IV discusses details of composition analysis. Structural characteristics of the mixed salts under consideration obtained within 8 Å around the central atom are presented in Sec. V. Results of molecular-dynamics simulations are discussed in Sec. VI. A general discussion of results is presented in Sec. VII and a summary and conclusion are presented in Sec. VIII.

II. EXPERIMENT AND SAMPLE PREPARATION

The reasons for our choice of the systems RbKBr and RbBrCl are as follows. First, two (Rb, Br) of the three constituent elements of the mixtures have convenient K edge energies (E0 = 15240 eV and 13500 eV, respectively) for XAFS measurements. Second, both systems are fully soluble in their solid states as shown in their phase diagrams with the minimal points of congruent melting where liquid and solid are in equilibrium at one concentration at 24% KBr in the first and 38% RbCl in the second mixture. This is the only assured condition to get a homogeneous solution in the bulk solid phase after freezing from the liquid, without additional annealing procedures. Any other composition typically crystallizes over a range of concentrations defined by the phase diagram and a cooling rate. Third, both systems have their consolute points (where they may theoretically separate into two solid phases) at rather low temperature (120 K for the first and 119 K for the second mixture) where the rate of mutual diffusion within the mixtures is negligibly small and one should not expect a phase

separation occurring during the low temperature XAFS measurements. Indeed, our thermal cycling around these points revealed no trace of the irreproducibility expected for the phase separation.

As described in Ref. 14, high purity (better than 99.9%) powders of raw materials of RbBr, KBr, and RbCl were supplied by Alfa (Johnson & Matthey). Two samples of mechanically mixed powders of above mentioned concentrations (Rb0.76K0.24Br and RbBr0.62Cl0.38), corresponding to their respective minimum melting temperature compositions, were prepared from these raw materials. Each of them was carefully mixed, placed into a quartz ampule, heated to 120°C for at least 1 h under vacuum of 10−3 Torr (to remove any trace of moisture since these salts are highly hygroscopic), and sealed off. Then the sealed ampules with the samples were transferred to a furnace and heated to 750°C (higher than the melting points of all of the three pure components) for 48 h to homogenize the solutions. They were then quenched into cold water to solidification and annealed afterwards during 4 days at the temperature 600°C (below the minimal melting points of the mixtures). The resultant mixed salts were bulk crystals with a lot of cracks and boundary lines. Finally these samples were removed from the quartz containers, ground to fine powders, sieved through 400 mesh, and rubbed as single layers onto Scotch tape and then sealed as a stacked array of layers in a vacuum tight cell, all within a dry glovebox.

To avoid a sample thickness effect the following condition should be satisfied for concentrated materials:

\[ \Delta \mu d \ll 1, \quad \Delta \mu z < 1.5, \]

where \( \Delta \mu \) is the absorption edge step, \( d \) is the average grain size, and \( z \) is the sample thickness. The thicknesses corresponding to \( \Delta \mu z = 1 \) for Rb0.76K0.24Br and RbBr0.62Cl0.38 at the Rb (Br) K edge are 67 (48) and 56 (78) \( \mu m \), respectively. To obtain samples with total \( \Delta \mu z \approx 1 \) required stacking at least 14 layers of the powdered tape. Thus the average grain size of the salts (\( d \approx 4 \mu m \)) was small enough to assure a negligible thickness effect.

The XAFS measurements were performed on the beam line X-11A at the National Synchrotron Light Source (NSLS) with a double-crystal (111) silicon monochromator. Harmonics were attenuated to negligible levels by detuning the monochromator to 80% of its peak. Due to the large thermal disorder of the investigated materials, the measurements had to be performed at low temperature and only the 30 K and 125 K data that had been measured were analyzed.

III. DATA ANALYSIS

The experimental XAFS data on mixed salts Rb0.76K0.24Br and RbBr0.62Cl0.38 at the congruent melting minimum points were analyzed in detail. The method of Ref. 17 was used to subtract the background function and obtain a normalized absorption function \( \chi(k) \):
FIG. 1. (a) Absorption coefficient $\mu(E)x$, and (b) normalized absorption $\chi(k)$ at 30 K of two independent measurements (dashed and solid lines) of the Br edge of Rb$_{0.76}$K$_{0.24}$Br.

$$
\chi(k) = \frac{\mu(k) - \mu_0(k)}{\Delta \mu_0(0)},
$$

where $k = \sqrt{2m(E - E_0)}$ is the wave number, $E$ is the photon energy, $\mu(k)$ is the measured absorption, $\mu_0(k)$ is the background, and $\Delta \mu_0(0)$ is the edge jump. The $\Delta \mu(E)x$ data and resultant $\chi(k)$ of two independent measurements of Rb$_{0.76}$K$_{0.24}$Br (Br edge) are shown in Fig. 1. The XAFS data were then fit to theoretical calculations, as discussed below. The fits were done in $R$ space, allowing shells higher than the first with their attendant multiple scattering to be correctly included in an organized way.

The reliability of multiple-scattering (MS) XAFS theory, the computer code FEFFS, and the fitting procedure used has been demonstrated before for the pure constituent compounds RbCl, KBr, and RbBr. In the pure salts, it was possible to obtain a fit up to 10 Å from the center atoms. However, to do so it was necessary to compensate for approximations in the theory by adding a small $\Delta E_0$ shift to the potential seen by near neighbors.

In this work the structure of mixed salts was determined using the same analysis procedure and the same scattering paths found to be important for XAFS of the pure salts RbBr, RbCl, and KBr. Because the structures of the mixed salts are more disordered, the analyses could not be extended past 8.5 Å for the ordered sublattice atoms, namely, the Br edge of Rb$_{0.76}$K$_{0.24}$Br and the Rb edge of RbBr$_{0.92}$Cl$_{0.08}$. For the disordered sublattice atoms, namely, the Rb edge of Rb$_{0.76}$K$_{0.24}$Br and the Br edge of RbBr$_{0.92}$Cl$_{0.08}$, the structure could not be reliably determined past 5.5 Å. Use of XAFS to investigate the structure of mixed salts was done previously, but only for the first nearest neighbor distributions.

The number of parameters $P$ which may be determined in XAFS analysis is limited to be less than the number of independent data points in the spectrum, $N$, given by

$$
N = \frac{2}{\pi} \delta k \delta R + 2,
$$

where $\delta k$ and $\delta R$ are, respectively, the ranges in $k$ and $R$ space over which the data are fit. The relevant quantities to determine $N$, such as $\delta k$, $\delta R$, and the Fourier transform window margins in $k$ space over which the data range $\delta k$ is gradually decreased to zero, the resultant $N$, and the subsequent $P$ values for all investigated mixtures are listed in Table I as is the $k^w$ factor which $\chi(k)$ is weighted with when fitting.

To best fit the data, three parameters for each important scattering path in the theory were allowed to vary. These adjustable parameters for each path were (1) $\Delta E_0$ (shift of the photoelectron energy origin), (2) $\Delta R$ (the shift of the atom from its site in the pure salt), and (3) $\sigma^2$ (Debye-Waller factor or second cumulant). An additional parameter, $\delta^2$ (the passive electron reduction factor for the central atom), was fixed to be the same as that found for the pure salt for each of the two central atoms, namely, 0.98 for Rb and 0.91 for Br. Because of geometric restrictions between scattering paths, some constraints could be placed on the variables, reducing the number of free parameters, $P$, in the fit.

Uncertainties in the $P$ free variables were determined by requiring the statistic $\Delta X^2$ for each fit to increase by 1 from the best-fit value, where $\Delta X^2$ is defined as

$$
\Delta X^2 = \frac{N}{n} \sum_{i=1}^{n} \left( \frac{\chi_{\text{data}}(R_i) - \chi_{\text{theory}}(R_i)}{\delta_i} \right)^2.
$$

<table>
<thead>
<tr>
<th>Data</th>
<th>Edge</th>
<th>$\delta k$ (Å$^{-1}$)</th>
<th>$\delta R$ (Å)</th>
<th>$k^w$</th>
<th>$dk_1$ (Å$^{-1}$)</th>
<th>$dk_2$ (Å$^{-1}$)</th>
<th>$N$</th>
<th>$P$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rb$<em>{0.76}$K$</em>{0.24}$Br</td>
<td>Br</td>
<td>3–11</td>
<td>2–8.5</td>
<td>$k$</td>
<td>1.8</td>
<td>3</td>
<td>34</td>
<td>24</td>
</tr>
<tr>
<td>Rb$<em>{0.76}$K$</em>{0.24}$Br</td>
<td>Rb</td>
<td>3–10</td>
<td>2–5.5</td>
<td>$k$</td>
<td>1.5</td>
<td>3</td>
<td>16</td>
<td>9</td>
</tr>
<tr>
<td>RbBr$<em>{0.92}$Cl$</em>{0.08}$</td>
<td>Br</td>
<td>2.5–10</td>
<td>2–5.5</td>
<td>$k$</td>
<td>1.2</td>
<td>3</td>
<td>18</td>
<td>9</td>
</tr>
<tr>
<td>RbBr$<em>{0.92}$Cl$</em>{0.08}$</td>
<td>Rb</td>
<td>2.5–11</td>
<td>2–8.5</td>
<td>$k$</td>
<td>1.5</td>
<td>1.5</td>
<td>36</td>
<td>25</td>
</tr>
</tbody>
</table>
Here $N$ is the number of independent data points [Eq. (4)], $\chi(R_i)$ is the Fourier transform of $\chi(k)$ at the point $R_i$, $n$ is the number of points used to evaluate $\chi(R_i)$ (including both real and imaginary parts) over the range $\delta R$, and $\delta_i^2$ is the sum of squares of the measurement uncertainty in $\chi_{\text{data}}(R_i)$ and the systematic uncertainties. In the cases considered here, the measurement uncertainty was very small and $\delta_i$ was determined mainly by systematic uncertainties of background subtraction and the approximations made in the theory of FEFF. The total systematic uncertainty was determined by requiring $\Delta \chi^2$ to be equal to $N - P$, as specified by statistical theory.\(^{21}\) Correlations between the fit variables were taken into account when evaluating the uncertainties, so that the uncertainty in a parameter is the value by which that parameter must be changed in order to increase $\Delta \chi^2$ by 1 when all other parameters are optimized. All uncertainties are indicated in the tables in parentheses.

### IV. COMPOSITION AND BOND LENGTH ANALYSIS

Consider the mixture $A_xB_{1-x}C$ of the salts $AC$ and $BC$ with composition $x$, and choose $C$ to be an absorbing atom. One has to perform a fit of the theoretical $\chi(k)$ to the unknown structure data. X-ray diffraction results show that the average structure of such a mixture remains a NaCl type and, hence, one may use for fitting only those paths which have been found important for the analysis of the pure components.\(^{14}\) The low $R$ region ($2 \, \text{Å} < r < 5.5 \, \text{Å}$) is dominated by contributions from the two nearest shells. It includes three kinds of single-scattering (SS) paths, two corresponding to the two different first nearest neighbor (1NN) $A$ or $B$ atoms ($c \rightarrow 1 \text{NN} \rightarrow c$, where $c$ is the center atom, which is a $C$ atom in this case) and one corresponding to the second nearest neighbor (2NN) distances $C-C$ ($c \rightarrow 2 \text{NN} \rightarrow c$).

The first shell XAFS, $\chi_{c,1}(k)$, is simply the average of the contributions of the two respective scatterings $\chi_{C_{AC}}(k)$ and $\chi_{C_{BC}}(k)$,

$$\chi_{1}(k) = y\chi_{C_{AC}}(k) + (1 - y)\chi_{C_{BC}}(k),$$  \hspace{1cm} (6)

where the composition factor $y$ is to be determined from the best fit of Eq. (6) to data. The resultant average $y$ value should be always $x$, regardless of the degree of randomness of the distribution of components since both the homogeneous mixture and heterogeneous separated structure (and any combination in between) should produce an XAFS signal averaged over the total number of absorbers $C$, with contributions $x$ and $1 - x$ of the components $CA$ and $CB$, respectively. The fitting of the 1NN paths to data was performed in accordance with Eq. (6). Indeed, the value of $y$ found agreed with $x$ within uncertainties (Table II) for both samples. It was also found that the $A-C$ and $B-C$ bond lengths $R_{\text{mix}}$ in both mixtures (Tables III, IV) did not follow Vegard's law [Eq. (1)]; i.e., they are not equal to one another and the VCA value, though sometimes the difference is barely outside of the uncertainties.

To find out the degree of randomness of the distribution realized in the mixture $A_xB_{1-x}C$ one needs to go to the 2NN and to investigate the $A$ or $B$ edge XAFS of this mixture. Now, if full phase separation occurs, the 2NN to an $A$ atom should be another $A$ atom with 100% probability. However, for the opposite case of a fully random mixture the second neighbor to an $A$ atom will be either again an $A$ or a $B$ atom in the ratio of $x$ to $1 - x$. Hence, it is straightforward to determine the degree of randomness from the $y$ value found from the best fit to data of $\chi_{2}(k)$ of the 2NN paths:

$$\chi_{2}(k) = y\chi_{AAA}(k) + (1 - y)\chi_{ABA}(k),$$  \hspace{1cm} (7)

where $\chi_{AAA}$ is the contribution for $A$ being both center and 2NN atoms, and $\chi_{ABA}$ is the contribution for $A$ being the center and $B$ the 2NN. Our previous analysis of the pure materials showed that only the SS paths give significant contributions to the 2NN peak.

The mixed sublattice atoms were analyzed, namely, the Br edge data of $\text{RbBr}_x\text{K}_{0.24}\text{Br}$ and the Br edge data of $\text{RbBr}_0.62\text{Cl}_{0.38}$. The fitting of the 2NN paths $\text{Rb} \rightarrow \text{Rb} \rightarrow \text{Rb}$ and $\text{Rb} \rightarrow \text{K} \rightarrow \text{Rb}$ to the second peak of the $\text{RbBr}_0.76\text{K}_{0.24}\text{Br}$ (Br edge) data [Fig. 2(a) and Table II] gave $y = 0.76 \pm 0.01$, demonstrating that complete

### TABLE II. Comparison of concentration $x$ and composition factor $y$ as determined from fits to the 1NN of $C$ and to the 2NN of $A$ atoms in $A_xB_{1-x}C$. Superscripts indicate the coordination shell.

<table>
<thead>
<tr>
<th>$A_xB_{1-x}C$</th>
<th>Bond</th>
<th>$x$</th>
<th>$y$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\text{RbBr}<em>0.76\text{K}</em>{0.24}\text{Br}$</td>
<td>Br-Rb$^1$</td>
<td>0.76</td>
<td>0.76(1)</td>
</tr>
<tr>
<td></td>
<td>Rb-Rb$^2$</td>
<td>0.76</td>
<td>0.76(1)</td>
</tr>
<tr>
<td>$\text{RbBr}<em>0.62\text{Cl}</em>{0.38}$</td>
<td>Rb-Br$^1$</td>
<td>0.62</td>
<td>0.62(1)</td>
</tr>
<tr>
<td></td>
<td>Br-Br$^2$</td>
<td>0.62</td>
<td>0.61(1)</td>
</tr>
</tbody>
</table>

### TABLE III. Four nearest bond lengths (in Å) about Br edge in pure RbBr, KBr, and $\text{RbBr}_0.76\text{K}_{0.24}\text{Br}$, and VCA values (Ref. 1). Superscripts indicate the coordination shell. The row with brackets gives the distance averaged over composition.

<table>
<thead>
<tr>
<th></th>
<th>Br-K$^1$</th>
<th>Br-Rb$^1$</th>
<th>Br-K$^2$</th>
<th>Br-K$^3$</th>
<th>Br-Rb$^3$</th>
<th>Br-Br$^4$</th>
</tr>
</thead>
<tbody>
<tr>
<td>$\text{RbBr}$</td>
<td>—</td>
<td>3.410(10)</td>
<td>4.822(10)</td>
<td>—</td>
<td>5.906(10)</td>
<td>6.820(10)</td>
</tr>
<tr>
<td>$\text{KBr}$</td>
<td>3.270(10)</td>
<td>—</td>
<td>4.624(10)</td>
<td>5.664(10)</td>
<td>—</td>
<td>6.540(10)</td>
</tr>
<tr>
<td>$\text{RbBr}<em>0.76\text{K}</em>{0.24}\text{Br}$</td>
<td>3.361(15)</td>
<td>3.393(10)</td>
<td>4.761(10)</td>
<td>5.860(20)</td>
<td>5.890(20)</td>
<td>6.743(15)</td>
</tr>
<tr>
<td>($\text{RbBr}<em>0.76\text{K}</em>{0.24}\text{Br}$)</td>
<td>3.385(11)</td>
<td>3.385(11)</td>
<td>4.761(10)</td>
<td>5.883(20)</td>
<td>5.883(20)</td>
<td>6.743(15)</td>
</tr>
</tbody>
</table>
TABLE IV. Four nearest bond lengths (in Å) about Rb edge in pure RbBr, RbCl, and RbBr0.62Cl0.38, and VCA values (Ref. 1). Superscripts indicate the coordination shell. The row with the brackets gives the distance averaged over composition.

<table>
<thead>
<tr>
<th></th>
<th>Rb-Cl(^1)</th>
<th>Rb-Br(^1)</th>
<th>Rb-Br(^2)</th>
<th>Rb-Cl(^1)</th>
<th>Rb-Br(^2)</th>
<th>Rb-Br(^4)</th>
</tr>
</thead>
<tbody>
<tr>
<td>RbBr</td>
<td>—</td>
<td>3.410(10)</td>
<td>4.822(10)</td>
<td>—</td>
<td>5.906(10)</td>
<td>6.820(10)</td>
</tr>
<tr>
<td>RbCl</td>
<td>3.270(10)</td>
<td>—</td>
<td>4.624(10)</td>
<td>5.664(10)</td>
<td>—</td>
<td>6.540(10)</td>
</tr>
<tr>
<td>RbBr0.62Cl0.38</td>
<td>3.336(15)</td>
<td>3.383(10)</td>
<td>4.753(10)</td>
<td>5.780(20)</td>
<td>5.810(20)</td>
<td>6.710(15)</td>
</tr>
<tr>
<td>(RbBr0.62Cl0.38)</td>
<td>3.305(12)</td>
<td>3.365(12)</td>
<td>4.753(10)</td>
<td>5.791(20)</td>
<td>5.791(20)</td>
<td>6.710(15)</td>
</tr>
</tbody>
</table>

random solubility occurred in the mixture preparation and therefore our following results in the larger R range, which assume this randomness, are trustworthy.

The first peak in R space of the Rb0.78K0.24Br X function [Fig. 2(a)] was fit by the single-scattering INN path Br-Br-Rb with the half path length 3.39 Å which is only 0.02 ± 0.015 Å shorter than that of pure RbBr (Table V). A similar analysis about the Br edge gave the same value for the Br-Rb nearest neighbor distance (Table III), as it should. This shortening of the first peak suggests that the first nearest neighbors bond length of pure salt may not be conserved in the mixture, in agreement with the results obtained in Refs. 3-7, but the difference for Br-Rb is too close to the uncertainty to be definitive. However, the analysis of the low-R-range data for the Br edge of the same mixed salt determined the Br-K 1NN distance to be 0.991 ± 0.015 Å longer than in pure KBr (Table III), distinctively different from the Br-Rb bond length change. Thus a first neighbor distance changes asymmetrically with concentration.

The same analysis was performed for the Br edge data of the other investigated mixture, RbBr0.62Cl0.38 [Fig. 2(b) and Table II] and the composition parameter of the components RbBr and RbCl was again determined from the fit to the second peak in R space by the 2NN SS paths Br → Br → Br and Br → Cl → Br. The obtained value γ = 0.61 ± 0.01 shows again that the components in the minimal melting point are randomly mixed with the desired concentration and the 1NN Rb-Br bond length (3.383 Å) also exhibited a small shift (∆R = -0.027±0.010 Å) relative to the pure RbBr bond length (Table VI). The same value of this bond length was obtained by analyzing the Rb edge data (Table IV). Here again the shift is still close to uncertainty to be definitive. However, analysis of the first peak in R space of the Rb edge for the same mixed salt gives a 0.06 ± 0.015 Å shift towards larger R in the Rb-Cl 1NN distance with respect to the pure RbCl (Table IV). Thus the shorter bond again changes dramatically in the mixture.

In the fits above we varied the shift of the photoelectron energy origin ΔE0 for each of the three considered paths independently (one 1NN and two 2NN SS paths for the mixed atom as the center atom, and two 1NN and one 2NN SS paths for the common atom as the center one). In the previous work14 on the pure salts it was shown that the paths containing first nearest neighbors must have an additional ΔE shift to the ΔE0 overall of all the other SS paths, which was interpreted as being caused by the lack of complete shielding of the core hole potential at the 1NN. The mixed salts indicate that a similar correction to FEFF5 is required for them also.

Analyzing the C-C (2NN) distances we found that in both Rb0.78K0.24Br and RbBr0.62Cl0.38 Vegard's law was valid within uncertainties for the lengths of Br-Br and Rb-Rb separations, respectively (Tables III, IV). These

FIG. 2. χ(R) of (a) Rb edge data of Rb0.78K0.24Br (solid line), pure RbBr (dotted line), and fit to the mixture (dashed line); (b) Br edge data of RbBr0.62Cl0.38 (solid line), pure RbBr (dotted line), and fit to the mixture (dashed line). The range over which the fit has been made is shown by the arrows. All the data are taken at T = 30 K.

TABLE V. Two nearest bond lengths (in Å) about Rb edge in pure RbBr, and Rb0.78K0.24Br, and VCA values (Ref. 1). Superscripts indicate the coordination shell. The row with brackets gives the distance averaged over composition.

<table>
<thead>
<tr>
<th></th>
<th>Rb-Br(^1)</th>
<th>Rb-K(^2)</th>
<th>Rb-Br(^2)</th>
</tr>
</thead>
<tbody>
<tr>
<td>RbBr</td>
<td>3.410(10)</td>
<td>—</td>
<td>4.822(10)</td>
</tr>
<tr>
<td>Rb0.78K0.24Br</td>
<td>3.390(10)</td>
<td>4.800(15)</td>
<td>4.790(15)</td>
</tr>
<tr>
<td>(Rb0.78K0.24Br)</td>
<td>3.390(10)</td>
<td>4.792(15)</td>
<td>4.792(15)</td>
</tr>
<tr>
<td>VCA</td>
<td>3.378</td>
<td>4.777</td>
<td>4.777</td>
</tr>
</tbody>
</table>
distances are identical to lattice constants defined by x-ray diffraction for these salts and do closely follow Vegard’s law. The violation of Vegard’s law for the 1NN bonds and its concurrent validity for the 2NN bonds lead to buckling of the bonds as discussed in Sec. V.

V. MULTIPLE SCATTERINGS AND DEVIATIONS FROM COLLINEARITY

Once the homogeneity and randomness of the mixtures \( A_xB_{1-x}C \) are verified, one can analyze the structure in the more distant \( R \) range which includes multiple scatterings. Our analysis was performed with the \( C \) edge data in both mixtures. This is the first time to our knowledge that the analysis of mixed salts beyond the first shell has been undertaken with an accurate account of multiple-scattering effects.

To extend the fitting procedure to the more distant \( R \) range, starting with the third nearest neighbor (3NN) distance, one should properly treat the single-scattering (SS) and multiple-scattering (MS) paths, contributing to this range. The variables should be carefully chosen in the fitting procedure so as to limit their number to the minimum required. Thus, we need to decrease the number of paths involved to only those that give significant contributions and to account for all correlations between the various parameters. As shown in Ref. 14 the most important paths needed for the analysis of salts with NaCl structure are single-scattering (SS), double-scattering (DS), and triple-scattering (TS) collinear paths. We, therefore, in the present work deal with such paths only.

The shortest paths beyond 2NN are SS paths \( c \rightarrow 3NN \rightarrow c \), corresponding to the pairs Br-Rb, Br-K for the Br edge of Rb0.76K0.24Br data and Rb-Br, Rb-Cl for the Rb edge of RbBr0.62Cl0.38 data. There are two fit variables assigned to each of the paths: \( \Delta R \) and \( \sigma^2 \). We fix the energy origin shift of these paths to be equal to the \( \Delta E_0 \) overall found for the 2NN bond Br-Br (Rb-Rb) and the composition parameter \( y = x \) as found in the previous section.

The next SS path connects the absorber and the fourth nearest neighbor (4NN). Again, we varied \( \Delta R \) as well as \( \sigma^2 \). There are four more paths to the 4NN in our analysis with the same half path length as the SS path which were fit to the data simultaneously since all of them contribute to the same \( R \) range 5–7 Å. They are double (DS) and triple-scattering (TS) focusing paths.

\[
\begin{array}{|c|c|c|}
\hline
\text{Br-Rb}^1 & \text{Br-Cl}^2 & \text{Br-Br}^2 \\
\hline
\text{RbBr} & 3.410(10) & 4.822(10) \\
\text{RbBr0.62Cl0.38} & 3.390(10) & 4.710(15) & 4.710(15) \\
\text{(RbBr0.62Cl0.38)} & 3.357 & 4.747 & 4.747 \\
\text{VCA} & & & \\
\hline
\end{array}
\]

\( c \rightarrow 1NN \rightarrow 4NN \rightarrow c \) (and its time reverse) and \( c \rightarrow 1NN \rightarrow 4NN \rightarrow 1NN \rightarrow c \), respectively. Each of these paths has two different contributions from the two different 1NN intervening atoms. Using the notation \( A_xB_{1-x}C \) for the mixed salts, we can denote these paths as follows. Absorber is \( C \), backscatter \( C' \), focusing atom \( A: C \rightarrow A \rightarrow C' \rightarrow C \) (DS) and its time reverse, and \( C \rightarrow A \rightarrow C' \rightarrow A \rightarrow C \) (TS); focusing atom \( B: C \rightarrow B \rightarrow C' \rightarrow C \) (DS), and its time reverse, and \( C \rightarrow B \rightarrow C' \rightarrow B \rightarrow C \) (TS). In these cases the \( C' \) atom is a 4NN and the focusing atom is a 1NN. Figure 3 represents schematically examples of SS, DS, and TS paths. The DS and TS paths are called focusing because the forward scattering of the intervening atom enhances the total signal. There are two more kinds of SS paths to the 5NN and 6NN which contribute to the total \( \chi(R) \) within \( R \) range 7–8 Å and were fit to data but they do not interfere with the shorter paths.

Since it was found from the fitting results of the 1NN paths that bond lengths \( C-A \) and \( C-B \) are different, the \( A \) and \( B \) atoms must be off their average lattice sites, because in the average structure these two bond lengths are equal. It can be shown that this means that the DS and TS paths are no longer collinear, contrary to what they are in the perfect structure. Describing the deviation from collinearity of these paths in terms of buckling angles \( \Theta_1 = 180^\circ - \angle ABC \) and \( \Theta_2 = 180^\circ - \angle ACB \), one can add these two more parameters (\( \Theta_1 \) and \( \Theta_2 \)) to focusing DS and TS paths and determine them from the fit.

A theoretical expression of \( \chi(k) \) of the MS path contains a product of scattering amplitudes \( F_i \), where \( i \) denotes the atomic site. The forward-scattering amplitude \( F \) of the intervening (focusing) atom in these paths as a function of \( \Theta \) has its maximum at collinearity (at \( \Theta = 0 \)) (see Ref. 22), and since it is even in \( \Theta \), an expansion

\[
\chi(k) = \sum_i F_i \sin(i\Theta)
\]

FIG. 3. Schematic of photoelectron SS(1), SS(2), SS(4), and SS(5) paths in the (100) plane of a NaCl structure to the 1NN, 2NN, 4NN, and 5NN, respectively, and DS (and its time reversed path DS) and TS paths to the 4NN. In all the cases the center atom is a positive ion denoted by c. SS paths to 3NN and 6NN are out of the plane.
of $F(k, \Theta)$ about $\Theta = 0$ has its lowest order of $\Theta^2$. Averaging over the total number of absorbing atoms, one obtains

$$
(F(k, \Theta)) \approx F(k, 0)[1 - b(k)(\Theta^2)]
$$

(8)

where the precisely forward-scattering amplitude of the intervening atom $F(k, 0)$ is determined with that FEFF5 calculation which gives good agreement with the ordered structure of the corresponding pure salts. The curvature coefficient $b(k)$ is obtained as follows. The $F(k, \Theta)$ is calculated with FEFF5 for the DS path described above with the focusing atom (the 1NN atom) moved out of its collinear site with varying $\Theta$ values. The $b(k)$ is then found by a fit to Eq. (8) in the small $\Theta$ range (up to $20^\circ$). The behavior of $F(k, \Theta)$ of the DS paths with Rb or Br as absorbers and Br, Cl or Rb, K as focusing atoms, respectively, for $k = 8 \text{ Å}^{-1}$ is shown on Fig. 4. Figure 5 shows the calculated $b(k)$ dependence for these paths. Since the Fourier transform window range $k_{\text{min}} = 2.5 \text{ Å}^{-1}$, $k_{\text{max}} = 11 \text{ Å}^{-1}$ with margins $dk = 1.5 \text{ Å}^{-1}$ (Table I) correspond to the approximately flat region of $b(k) \approx 0.0018 \text{ deg}^{-2}$, one can fix $b(k)$ to that constant and introduce the new fit variable $(\Theta^2)$.


![FIG. 5. Curvature coefficient $b(k)$ for the DS paths. Absorber Br with focusing atoms K (solid line) and Rb (dashed line), and absorber Rb with focusing atoms Br (dotted line) and Cl (long dashed line).](image)

![FIG. 7. Fit of the theoretical $\chi(R)$ (dashed line) to the Br edge data of Rb$_{0.76}$K$_{0.24}$Br (solid line) at (a) 30 K and (b) 125 K. The range over which the fit has been made is shown by the arrows.](image)
FIG. 8. Fit of the theoretical $\chi(R)$ (dashed line) to the Rb edge data of RbBr$_{0.62}$Cl$_{0.38}$ (solid line) at (a) 30 K and (b) 125 K. The range over which the fit has been made is shown by the arrows.

The fit was performed in the $R$ range 5–7 Å for both mixtures. The paths involved were the above described two kinds of SS paths (c → 3NN → c and c → 4NN → c), two kinds of DS and TS paths. Figure 6 gives the contributions of different paths to the final fit in the corresponding $R$ range. These DS and TS contributions are not small and therefore give us a real possibility to measure buckling angles.

Note that the TS path requires only one fitting variable additional to the DS paths, namely, a $\Delta E_0$ shift: The focusing atom (1NN) scatters the photoelectron twice and the amplitude of $\chi(k)$ is multiplied by the factor $(1 - 0.0018(\Theta^2))^2$.

Another fit variable, $\Delta R$ of the TS path, differs generally from that of the corresponding DS path. However, in our case, where the difference in bond lengths AC and BC is relatively small, we expect the deviations from the average structure to be also small and the rms bond angles $(\Theta^2)^{1/2}$ not to exceed 20°. Thus, the difference in the half path length of the DS and TS paths is expected not to exceed 0.015 Å, which is the uncertainty of the distance determination with XAFS (see Tables III, IV) and one can set $\Delta R_{DS} = \Delta R_{TS}$ to a good approximation. This also implies that the Debye-Waller factors of the DS and TS paths, connecting the central atom, focusing atom (1NN) and the 4NN atom are approximately equal to the $\sigma^2$ of the single pair c → 4NN due to the relation $\sigma^2_{DS} = \sigma^2_{TS} = \sigma^2_{c4NN}$, verified to be valid in the analysis of pure salts.

The fit results up to $R = 8$ Å for both mixtures at 30 K are shown on Figs. 7(a) and 8(a). As a final check of our approximation of a constant $b$, we set the $(\Theta^2)$ values to those found in our fit and then calculated the XAFS spectrum using the $b(k)$ variable in $k$ and obtained fits which were indistinguishable from the one obtained with the constant $b$. Figures 7(b) and 8(b) show fits to only the two nearest neighbors at 125 K (Rb$_{0.78}$K$_{0.24}$Br, Br edge, and RbBr$_{0.62}$Cl$_{0.38}$, Rb edge, respectively), in order to determine the 1NN temperature dependence of $\sigma^2$.

Values of fit variables including obtained bond angles are given in Tables VII, VIII. The rms deviations from collinearity actually are less than 10°, validating the approximations used.

<table>
<thead>
<tr>
<th>Path</th>
<th>$\Delta E_0$ (eV)</th>
<th>$\sigma^2_{min}$</th>
<th>$\sigma^2_{Br}$</th>
<th>$\sigma^2_{KB}$</th>
<th>$\Theta$ (deg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Br → K$^+$ → Br</td>
<td>2.6(5)</td>
<td>0.64(5)</td>
<td></td>
<td>0.61(5)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>[1.4(1)]</td>
<td>[1.0(1)]</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Br → Rb$^+$ → Br</td>
<td>3.6(5)</td>
<td>0.60(5)</td>
<td>0.54(5)</td>
<td></td>
<td></td>
</tr>
<tr>
<td></td>
<td>[1.1(1)]</td>
<td>[1.06(5)]</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Br → Br$^+$ → Br</td>
<td>0.8(5)</td>
<td>1.30(5)</td>
<td>0.73(5)</td>
<td>0.51(5)</td>
<td></td>
</tr>
<tr>
<td></td>
<td>[1.1(1)]</td>
<td>[1.06(5)]</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Br → K$^+$ → Br</td>
<td>0.8(5)</td>
<td>1.62(5)</td>
<td></td>
<td>0.87(5)</td>
<td></td>
</tr>
<tr>
<td>Br → Br$^+$ → Br</td>
<td>0.8(5)</td>
<td>1.36(5)</td>
<td>1.11(1)</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Br → Br$^+$ → Br</td>
<td>0.8(5)</td>
<td>1.00(5)</td>
<td>0.77(5)</td>
<td>0.57(5)</td>
<td></td>
</tr>
<tr>
<td>Br → K$^+$ → Br</td>
<td>0.0(5)</td>
<td>1.00(5)</td>
<td></td>
<td>0.57(5)</td>
<td>7.7(2.7)$^\circ$</td>
</tr>
<tr>
<td>Br → Rb$^+$ → Br</td>
<td>0.0(5)</td>
<td>1.00(5)</td>
<td>0.77(5)</td>
<td></td>
<td>9.0(1.6)$^\circ$</td>
</tr>
<tr>
<td>Br → K$^+$ → Br</td>
<td>-0.5(5)</td>
<td>1.00(5)</td>
<td>0.57(5)</td>
<td></td>
<td>7.7(2.7)$^\circ$</td>
</tr>
<tr>
<td>Br → Rb$^+$ → Br</td>
<td>-0.5(5)</td>
<td>1.00(5)</td>
<td></td>
<td></td>
<td>9.0(1.6)$^\circ$</td>
</tr>
<tr>
<td>Br → K$^+$ → Br</td>
<td>0.8(5)</td>
<td>1.21(1)</td>
<td></td>
<td>0.85(5)</td>
<td></td>
</tr>
<tr>
<td>Br → Rb$^+$ → Br</td>
<td>0.8(5)</td>
<td>1.4(1)</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Br → Br$^+$ → Br</td>
<td>0.8(5)</td>
<td>1.45(10)</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
VI. COMPUTER SIMULATION

Using the XAFS results for the average values, a model of the structure showing the distribution of bond lengths and bond angles of the mixed ionic salts under consideration may be calculated using the molecular-dynamics (MD) simulation method, in which the equations of lattice motion are integrated numerically. The atoms in ionic salts, especially alkali halides, have a high degree of charge transfer and their interatomic potentials consist of a sum of Coulomb and repulsion terms. The anharmonicity of the atomic vibrations are found to become significant only well above 30 K, and so at 30 K these potentials $V$ may be approximated as harmonic,

$$V = \frac{1}{2} \sum_{i<j} k_{ij} (R_{ij} - \rho_{ij})^2,$$

where $k_{ij}$ is the effective force constant, and $R_{ij}$ and $\rho_{ij}$ are the distances between the $i$ and $j$ ions and their equilibrium bond lengths, respectively. We add the approximation in Eq. (9) that the interactions are significant only between the first nearest neighbors. Therefore, in the case of the NaCl-type cubic lattice the summation is performed over the six nearest neighbor bonds.

We used in this work the simulation method$^{23}$ and computer code CLUSTER$^{23}$ which showed good agreement with experiment for the simulation of pure alkali halides. The equations of motions were integrated in Ref. 23 using a fifth-order predictor-corrector algorithm. The program CLUSTER was applied in our work to calculate the mixed salts with the interatomic potentials of Eq. (9), where $i$ and $j$ are the nearest neighbors, $\rho_{ij} = \rho_{AC}$ or $\rho_{BC}$, the average lengths of the bonds $A-C$ or $B-C$, respectively, as determined here by XAFS, and $k_{AC}$ and $k_{BC}$ are the force constants of these bonds.

At low temperature, where zero point motion dominates, the force constants may be obtained from the Einstein formula$^{25}$ for the Debye-Waller factor:

$$\sigma^2 \approx \frac{\hbar}{2\mu \omega},$$

where $\omega$ is the oscillation frequency of the pair and $\mu = (M_i^{-1} + M_j^{-1})^{-1}$ is its reduced mass. Finding $\omega$ from the values of $\sigma^2$ of the first nearest neighbor bonds, determined for pure salts,$^{14}$ one can calculate $k_{ij} = \mu \omega^2$.

We assume that the $k_{ij}$ determined from the pure salts are a good enough approximation for the mixed salt case. Our result is not sensitive to this approximation. Table IX contains the values of $\sigma^2$, $\mu$, $\omega$, and $k_{ij}$ calculated for the Rb-Br, Rb-Cl, and K-Br bonds in the pure salts.

The simulation was performed over both $9 \times 9 \times 9$ (729 atoms) and $11 \times 11 \times 11$ (1331 atoms) clusters in order to check for surface effects. The initial configuration $A_n B_{1-n} C$ was prepared using a random number generator for placing $A$ and $B$ atoms at their sites in accordance with the concentration $x$. Atoms $C$ were placed at the sites of the average lattice with the lattice parameter $a = 6.74$ Å for Br-Br (4NN) distance in Rb$_{7.66}$K$_{0.24}$Br and $a = 6.71$ Å for Rb-Br (4NN) distance in RbBr$_{0.62}$Cl$_{0.38}$, as determined from our XAFS analysis. The atoms $A$ and $B$ were initially placed in the middle of the $C$-C bond. In order to satisfy the fit results for the first path (Tables III, IV) we set $\rho_{AC} = \rho_{RB-Br} = 3.39$ Å, $\rho_{BC} = \rho_{K-Br} = 3.36$ Å in Rb$_{7.66}$K$_{0.24}$Br, $\rho_{AC} = \rho_{RB-Br} = 3.38$ Å, $\rho_{BC} = \rho_{Rb-Cl} = 3.34$ Å in RbBr$_{0.62}$Cl$_{0.38}$.

During the subsequent MD run the surface atoms were

\[ \text{TABLE VIII. Path parameters $\Delta E_i$ (in eV) and $\sigma^2_i$ (in $10^{-2}$ Å$^2$) of RbBr$_{0.62}$Cl$_{0.38}$ (Rb edge) data at 30 K. DW factors of the 1NN paths at 125 K are given in brackets. $\sigma^2_i$ of pure salts (Ref. 14) RbBr and RbCl (Rb edge) are given for comparison. Superscripts indicate the coordination shell.} \]

<table>
<thead>
<tr>
<th>Path</th>
<th>$\Delta E_i$</th>
<th>$\sigma^2_i$</th>
<th>$\sigma^2_{RbBr}$</th>
<th>$\sigma^2_{RbCl}$</th>
<th>$\Theta$ (deg)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rb$\rightarrow$Cl$^1$ $\rightarrow$ Rb</td>
<td>1.3(5)</td>
<td>0.96(5)</td>
<td>0.55(5)</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Rb$\rightarrow$Br$^1$ $\rightarrow$ Rb</td>
<td>-1.3(5)</td>
<td>0.60(5)</td>
<td>0.55(5)</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Rb$\rightarrow$Rb$^2$ $\rightarrow$ Rb</td>
<td>-2.4(5)</td>
<td>1.27(5)</td>
<td>0.71(5)</td>
<td>0.40(5)</td>
<td>7.3(3.0)$^*$</td>
</tr>
<tr>
<td>Rb$\rightarrow$Cl$^2$ $\rightarrow$ Rb</td>
<td>-2.4(5)</td>
<td>1.40(5)</td>
<td>0.55(5)</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Rb$\rightarrow$Br$^3$ $\rightarrow$ Rb</td>
<td>-2.4(5)</td>
<td>1.50(5)</td>
<td>1.06(5)</td>
<td>-</td>
<td>-</td>
</tr>
<tr>
<td>Rb$\rightarrow$Rb$^4$ $\rightarrow$ Rb</td>
<td>-2.4(5)</td>
<td>1.40(5)</td>
<td>0.81(5)</td>
<td>0.44(5)</td>
<td>8.2(1.5)$^*$</td>
</tr>
<tr>
<td>Rb$\rightarrow$Cl$^1$ $\rightarrow$ Rb$^4$ $\rightarrow$ Rb$^4$ $\rightarrow$ Rb</td>
<td>0.0(5)</td>
<td>1.04(5)</td>
<td>-</td>
<td>7.3(3.0)$^*$</td>
<td>8.2(1.5)$^*$</td>
</tr>
<tr>
<td>Rb$\rightarrow$Cl$^1$ $\rightarrow$ Rb$^4$ $\rightarrow$ Cl$^1$ $\rightarrow$ Rb</td>
<td>-2.0(5)</td>
<td>1.40(5)</td>
<td>-</td>
<td>0.44(5)</td>
<td>8.2(1.5)$^*$</td>
</tr>
<tr>
<td>Rb$\rightarrow$Br$^1$ $\rightarrow$ Rb$^4$ $\rightarrow$ Br$^1$ $\rightarrow$ Rb</td>
<td>-2.0(5)</td>
<td>1.40(5)</td>
<td>0.81(5)</td>
<td>-</td>
<td>8.2(1.5)$^*$</td>
</tr>
</tbody>
</table>

\[ \text{TABLE IX. $\sigma^2$, reduced masses, Einstein frequencies, and force constants of Rb-Br, K-Br and Rb-Cl bonds in our computer simulations.} \]

<table>
<thead>
<tr>
<th>Bond</th>
<th>$\sigma^2$ ($10^{-3}$ Å$^2$)</th>
<th>$\mu$ (10^{-27}$ kg$)</th>
<th>$\omega$ (10^{12}$ Hz$)</th>
<th>$k_{ij}$ (N/m)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rb-Br</td>
<td>5.5</td>
<td>68.6</td>
<td>1.4</td>
<td>13.4</td>
</tr>
<tr>
<td>K-Br</td>
<td>6.1</td>
<td>43.8</td>
<td>2.0</td>
<td>17.4</td>
</tr>
<tr>
<td>Rb-Cl</td>
<td>5.5</td>
<td>41.5</td>
<td>2.3</td>
<td>22.0</td>
</tr>
</tbody>
</table>
fixed by setting their velocities to zero. The internal atoms were allowed to move in accordance with equations of motion and initially given random velocities which were then rescaled several times in order to obtain a stable equilibrium configuration in the minimum of the potential well. The time step was chosen to be $10^{-15}$ sec [it must be at least an order of magnitude smaller than the bond oscillation period $\omega^{-1}$ (Table IX)]. For both clusters averaging was done only over the inner $5 \times 5 \times 5$ cluster to exclude the influence of the fixed surface atoms. For the larger cluster, averaging was also done over the inner $7 \times 7 \times 7$ cluster to verify that the averaged region had been immersed into the interior deeply enough to be insulated from surface effects. The time averaging ($2 \times 10^{-11}$ sec) was performed during the last run of the program where the desired equilibrium was achieved. The results for angles obtained from the various clusters agreed one with another to about $0.1^\circ$, indicating that surface effects were negligible.

The simulation results in equilibrium were found to be in excellent agreement with the XAFS results. The equilibrium bond lengths $\rho_{\text{Rb-Br}} = 3.39 \text{ Å}$, $\rho_{\text{K-Br}} = 3.37 \text{ Å}$, and $\rho_{\text{Rb-CI}} = 3.34 \text{ Å}$ are consistent with those determined by XAFS within the uncertainty $0.01 \text{ Å}$. The rms bond angles found with MD simulation for the final configuration of atoms were $\Theta(\text{Br-Rb-Br}) = 8.2^\circ$, $\Theta(\text{Br-K-Br}) = 8.0^\circ$ in Rb$_0.76$K$_0.24$Br and $\Theta(\text{Br-Br-Rb}) = 8.8^\circ$, $\Theta(\text{Br-CI-Rb}) = 8.0^\circ$ in RbBr$_0.62$Cl$_0.38$, agreeing within uncertainties with the measurements (Tables VII, VIII). Figures 9(a) and 9(b) show the equilibrium configurations of Rb$_0.76$K$_0.24$Br and RbBr$_0.62$Cl$_0.38$ clusters, respectively. The static distortions off NaCl structure sites are clearly seen on both plots. Within the uncertainties these pictures should be considered as actual structures of our mixed salts.

VII. DISCUSSION

Our XAFS results give microscopic local information on the structure of the two mixed salts Rb$_0.76$K$_0.24$Br and RbBr$_0.62$Cl$_0.38$ which can hardly be obtained by other means. The distances between pairs of atoms, the three-body correlations expressed in terms of the buckling angles from collinearity, the relative amplitudes of vibration of the atoms, and the evidence that their two component salts are randomly mixed were derived from XAFS. This microscopic local information determines the detailed differences between the actual structure and the average structure of mixed salts as ascertained by diffraction.

In this case the analysis could proceed to neighbors beyond the second since the the samples are known to be homogeneous and the distribution of the components is random. The analysis of the multiple scatterings between the three atoms comprising the center atom, a nearest neighbor and that fourth neighbor that would be collinear with the first two in the corresponding pure components, found that the three atoms are no longer collinear in the mixture. The XAFS determined the rms angular deviation from collinearity of $7.2\pm3.0^\circ$ and $8.2\pm1.5^\circ$ for the Rb-Cl-Rb and Rb-Br-Rb atoms, respectively, in RbBr$_0.62$Cl$_0.38$. Similarly, the XAFS determined the rms angular deviation from collinearity of $7.7\pm2.7^\circ$ and $9.0\pm1.6^\circ$ for the Br-K-Br and Br-Br-Br atoms, respectively, in Rb$_0.76$K$_0.24$Br. This buckling is consistent with both the XAFS-determined deviations of the nearest neighbor distances from the average structure ones and the nearest neighbor force constants between the various pairs of atoms, as shown by the molecular-dynamics simulation. The simulation graphically illustrates the local structure and how it deviates from the average in Fig. 9. In fact, this represents a determination of the actual structure as opposed to the periodic average structure determined from diffraction.

It should be emphasized that the determination of the buckling is a direct measurement by XAFS of three-body correlations, and it is only possible with the high accuracy shown here because of very strong multiple scatterings and their high sensitivity to the small deviations from collinearity in this special case of the NaCl-type average structure. Ordinary elastic $x$-ray or neutron scattering cannot directly determine three-body correlations, but are limited to only two-body correlations. The XAFS results also found that the distance between pairs of atoms and their mean square disorder $\sigma^2$ are different between the different component atoms, even at the same neighboring shell, a result which cannot be determined from standard diffraction. The XAFS measurements, together with the information from diffraction that the average structure is a NaCl type, give all the information necessary for a modeling, and, thus, for solving the local structure of the mixed salts by a molecular-dynamic simulation, as shown in Fig. 9.

The buckling could be inferred from the fact that twice the average 1NN distance (denoted in Tables III and IV by brackets enclosing the chemical formula) found by averaging over concentration the respective 1NN bond lengths determined by XAFS for each pair is larger by
0.03 ± 0.022 Å than the 4NN distance. However, this is not convincingly outside the uncertainties. On the other hand the directly measured buckling angle is clearly significant and convincingly proves the presence of buckling. Actually, it is sufficient that the 1NN distances of each component be different from the VCA value for buckling to occur, even if their average is still close to the VCA value. To prove this we have used the values of the 1NN distances presented by Boyce and Mikkelsen for the congruent melting composition of the Rb$_2$K$_{1-x}$Br mixed salt [Rb-Br = 3.389 Å, K-Br = 3.319 Å, whose compositional average is the VCA value (Fig. 10)] keeping all other distances the same in a molecular-dynamics simulation. We have found an rms Θ about 0.6° larger than the value found for our 1NN distances, but the same within experimental uncertainties. The change in rms Θ is small in this case because the majority atom bond Rb-Br did not change in the simulation, only the smaller concentration one, K-Br. When we changed the Rb-Br distance in our simulation, the rms Θ did change strongly. This test characterizes the different sensitivity of the rms buckling angle to the influence of different atoms in mixture, as well as double-checks our XAFS results.

Although almost no differences (within uncertainties) were observed between the buckling angles of the both systems of mixed salts, the mixture RbBr$_{0.92}$Cl$_{0.08}$ can be characterized as stronger distorted than the second one, Rb$_{0.76}$K$_{0.24}$Br. Tables III–VI clearly show that greater deviations from the average structure occurred for the 1NN distributions in the RbBr$_{0.92}$Cl$_{0.08}$ mixture, in spite of the fact that the initial difference in the size of the two mixing atoms was the same for both systems. Interestingly, as one might expect, the deviations from the average structure in the $A_2B_{1-x}C$ mixture are correlated with the difference in the force constants of the two 1NN pairs: $A-C$ and $B-C$. The smaller the difference $k_{AC}-k_{BC}$, the smaller are the deviations of the atoms from their average sites. In our systems of mixed salts the difference between force constants of Rb-Br and Rb-Cl pairs is larger than that of Br-Br and K-Br (Table IX). Correspondingly, the mixture RbBr$_{0.92}$Cl$_{0.08}$ is found to be more distorted than the Rb$_{0.76}$K$_{0.24}$Br. This is understandable as the positively charged cations are small, tightly bound, while the negatively charged anions are comparatively large, loosely bound configurations of atoms. It takes more energy to distort the cations and thus to mold their size to the average, and it is expected that a disordered sublattice of cations will have more deviation from an average structure.

Our results clearly show that the concept of fixed size bonds in alkali halides as they are mixed is quite incorrect. Substantial changes occurred for the Rb-Cl (0.07 Å) and K-Br (0.09 Å) 1NN distances as their salts were mixed into a single phase. The other point to note from Tables III, IV, VII, and VIII is the comparison of these changes occurring in bond lengths with differences of their $\sigma^2$, mean squared disorder, between the mixed salts and the corresponding values of the pure salts. For instance, the Br-K 1NN distance changes from 3.27 Å in the pure salt to 3.36 Å in the mixed salt (Table III) with no significant increase in $\sigma^2$ (Table VII). This large change of 0.09 Å with no change in $\sigma^2$ indicates that the ions remain in contact and thus their size is concentration dependent. Just the fact that the bond distance changes is, by itself, insufficient evidence of a change in size of the ions because, as pointed out by Ashcroft and Mermin, the ions may be not “touching.” The evidence of touching, though, is reflected by the $\sigma^2$ since this is a measure, at the low temperatures of the measurements, of the zero point motion and thus the contact force between atoms. The lack of change in $\sigma^2$ indicates the same contact force and thus “touching” for both distances of this mixture. In contrast, the Rb-Cl 1NN distance increases in the mixed salt over the pure salt by 0.07 Å (Table IV). Yet the corresponding value of the pure salt $\sigma^2$ also almost doubles in the mixture (Table VIII). It suggests that the contact present in the pure salt is loosened in the mixture (and therefore the atoms are no longer “touching”) and the increase in bond length should not be interpreted as an increase in ionic size. To verify this conclusion we have studied the Debye-Waller factors of these bonds at 125 K. The results (Tables VII and VIII) show that the Rb-Cl $\sigma^2$ in the mixed salt has also a larger temperature dependence than the other 1NN bonds, which is consistent with the conclusion that the atoms are no longer “touching.”

First neighbor distances have been measured previously in the same mixed salt systems and Fig. 10 presents the result of comparison of Refs. 3, 6 with our data. We have a rather good agreement with published

![Graph](image-url)
data for the longer Rb-Br bond (which contracts under mixing) in both mixtures. However, for the shorter bond, K-Br, which expands under mixing, our result has a larger value than that of Boyce and Mikkelsen. Although this difference appears in Fig. 10(a) to be outside of the uncertainties of Ref. 3, a private communication with Boyce leads us to accept that the real uncertainties in his results may be large enough to cover the disagreement. The solid line in Fig. 10 is the VCA value and in order for the average of the first neighbor distances to agree with the VCA model the shorter bond needs to lie on the lower dashed lines. The results of Boyce and Mikkelsen as displayed in Fig. 10(a) led the authors to argue that the average of the first bond neighbors does satisfy the VCA model; thus a result that they implied was to be expected. However, when one realizes that the buckling does occur, there is no reason to expect the average of the two 1NN distances to be the VCA value, in agreement with what we find. Our results indicate that it is easier for a shorter bond to expand than for a longer one to contract. This fits the known fact of wider solubility limits of smaller atoms in the midst of larger ones (if other factors are similar) in mixed crystals, and also is as expected from the asymmetric form of the pair potential between atoms where the repulsive force is much larger than the attractive one. Another factor is that one expects that the negatively charged Br ion can be distorted more easily than the positively charged K core and thus it is likely that the negative ion has the more variable size.

In an interesting set of papers, Thorpe and collaborators have calculated the distribution of the disorder about the average zinc-blende or diamond structure for random semiconductor alloys, but did not use this information to try to solve the structure. They assumed that the alloys retain the same elastic constants as the pure constituents and obtain excellent agreement with experimental XAFS measurements of interatomic distances as a function of concentration in most cases. Our results, as discussed just above, show that this assumption of unmodified elastic constants is not valid for the mixed salts, which would lead to an incorrect conclusion about the behavior of the Rb-Cl bond in the mixture.

It should be emphasized that the changes revealed between the pure salts and their mixtures are reliably detected because the theory has been calibrated against the pure salts where the average and local structures are the same. In essence, we are depending on the accuracy of the theory to determine only the changes between the pure and mixed salts and a 10% error in the theory gives only a 10% error in the difference. Thus, the difference is known with greater accuracy than has been estimated in this paper where it was assumed that the errors in the total distance between atoms are uncorrelated between the pure and mixed salts. Only in the determination of \( \Theta_{\text{rms}} \), the rms buckling angle of the mixture, where the change is directly determined, are the errors not overestimated. It should also be pointed out that the accuracy of our method to determine \( \Theta_{\text{rms}} \) has been proven by testing against NaTaO$_3$ whose value of \( \Theta_{\text{rms}} \) is known at low temperatures from diffraction measurements.

The value of \( \Theta_{\text{rms}} \) determined by our measurements is a quadrature sum of contributions from both the buckling and zero point motion. We estimate zero point contribution from the \( \sigma^2 \) listed in Tables VII and VIII. The \( \Theta_{\text{rms}}^{\text{zpm}} \) due to zero point motion is \( 2\sigma_1/R_1 \) where \( \sigma_1 \) is an average 1NN value of zero point disorder and \( R_1 \) is an average 1NN distance, giving \( \Theta_{\text{rms}}^{\text{zpm}} = 2.6^\circ \). The factor of 2 comes from the definition of \( \Theta_{\text{rms}} \) which is twice the rms angle between the collinear direction and the 1NN bond. This vibrational contribution causes an increase in \( \Theta_{\text{rms}} \) of less than 7–8%, and thus is negligible since it is less than the uncertainties.

**VIII. SUMMARY AND CONCLUSIONS**

The XAFS measurements and data analysis were made of the mixed salts Rb$_x$K$_{1-x}$Br and RbBr$_x$Cl$_{1-x}$ at 30 K and 125 K. The concentration \( x \) of each mixture was chosen at the minimum congruent melting point. The XAFS results verified that the samples were homogeneous and randomly mixed. Thus it became possible to give a full detailed description of the difference between the actual structure of the mixtures and the average structure as measured by diffraction. The cations in the first mixture and anions in the second one were shown to be shifted from their average sites forming a buckled NaCl-type crystalline structure with an rms angular deviations from collinearity (defined as twice the angle between a 1NN bond and the collinear direction) of \( 7^\circ-9^\circ \). The distance between the nearest atoms deviated from the average values while the \( \sigma^2 \) disorder in these distances depended on the particular pair. A molecular-dynamics simulation using as input parameters the values determined from the XAFS measurements was able to reproduce the measured results and to give a detailed description of the actual structure of the mixture as shown in Fig. 9. Although diffraction determines that the average structure of these mixed salts is a NaCl one, the results presented above solve the actual structure and display the fluctuations about the average.

The deviations from the average are not simply dependent on the difference in sizes of the atoms that are being mixed. In fact, our results, in agreement with others, indicate that the size of an atom is not an accurate concept since it is not fixed. The Rb-Cl and K-Br 1NN distances change by 0.066 \( \pm \) 0.015 Å and 0.091 \( \pm \) 0.015 Å, respectively, between the corresponding pure and chosen mixed salts. In addition, the Rb-Cl bond in the mixture becomes much weakened suggesting that the atoms no longer touch, making it no longer meaningful to associate the bond to a size of these atoms which change with concentration. We find it more meaningful to consider both bond lengths and force constants. For example, the mixture RbBr$_x$Cl$_{1-x}$ with a higher difference in 1NN force constants exhibits larger distortions than the other one, Rb$_x$K$_{1-x}$Br.

The ability of XAFS to determine three-body correlations with high accuracy in the special case of near collinearity deserves a special mention. This feature had been appreciated for some time but the realization of
this potential had to await the development of an accurate theory such as that used in the computer code FEPS.\textsuperscript{13}

The results of this investigation show important differences between the average structure and the real structure of these mixtures. It is clear that any accurate theory of the properties of these mixtures must include these deviations. For example, it has been pointed out that the melting of disordered mixtures depends on deviations from the average structure. The more the disorder, the greater is the depression of the melting temperature.\textsuperscript{8,12} The range of solubility of two components typically depends on the differences in their atomic "sizes"; the greater their difference, the smaller the range. The electrical resistivity, dielectric, optical, and mechanical properties are expected to depend on this disorder as well. Accurate theoretical calculations of the properties of mixed salts and other disordered solids need to account for the deviations of the structure from the average. Our results now permit such calculations to be done since they determine the actual structure. Coordinates of the atomic positions of these salts obtained from MD simulations are available for such purposes.\textsuperscript{29}

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